Store at -20°C

#89052

SimpleChIP[®] Mouse TULP4 Promoter Primers

500 µl (250 PCR reactions)



Support: +1-978-867-2388 (U.S.) www.cellsignal.com/support

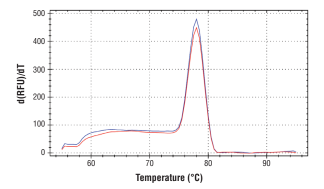
Orders: 877-616-2355 (U.S.) orders@cellsignal.com

Entrez-Gene ID #68842 UniProt ID #Q9JIL5

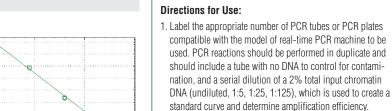
For Research Use Only. Not For Use In Diagnostic Procedures.

Applications	Species Cross-Reactivit	ty	Primer/Anneal Extensi	ion PCR Poduct Length
ChIP	M		65°C	124 bp
Description: SimpleChIP® Mouse TL contain a mix of forward and reverse P specific to a region of the mouse tubby promoter. These primers can be used t been isolated using chromatin immund Primers have been optimized for use in tive real-time PCR and have been testes SimpleChIP® Enzymatic Chromatin IP and ChIP-validated antibodies from Ce	CR primers that are ' like protein 4 (TULP4) o amplify DNA that has oprecipitation (ChIP). I SYBR [®] Green quantita- d in conjunction with Kits #9004 and #9005	Threshold Cycle (C_r)	31 30 29 28 27 26 25 24 -2 -1 Log Startin E= 105.2%	0 1.0 g Quantity (ng) R ² = 0.997

SimpleChIP® Mouse TULP4 Promoter Primers were tested on DNA isolated from cross-linked cells using the SimpleChIP® Plus Enzymatic Chromatin IP Kit (Magnetic Beads) #9005. Real-time PCR was performed in duplicate on a serial dilution of 2% total input DNA (20 ng, 4 ng, 0.8 ng, and 0.16 ng) using a real-time PCR detection system and SYBR® Green reaction mix. The PCR amplification efficiency (E) and correlation coefficient (R²) were calculated based on the corresponding threshold cycle (C₂) of each dilution sample during 40 cycles of real-time PCR (95°C denaturation for 15 sec, 65°C anneal/extension for 60 sec).



PCR product melting curves were obtained for real-time PCR reactions performed using SimpleChIP® Mouse TULP4 Promoter Primers. Data is shown for both duplicate PCR reactions using 20 ng of total DNA. The melt curve consists of 80 melt cycles, starting at 55°C with increments of 0.5°C per cycle. Each peak is formed from the degradation of a single PCR product.



2.0

at -20°C.

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 Add 2 µl of the appropriate ChIP DNA sample to each tube or well of the PCR plate.

Storage: Supplied in nuclease-free water at a concentration of

5 μ M (each primer is at a final concentration of 5 μ M). Store

 Prepare a master PCR reaction mix as described below. Add enough reagents for two extra reactions to account for loss of volume. Add 18 µl of the master PCR reaction mix to each PCR reaction tube or well of the PCR plate.

Reagent	Volume for 1 PCR Reaction (20 µl)
Nuclease-free H ₂ O		6 µl
5 μM SimpleChĺP®	Primers	2 µl
2X SYBR® Green Re	action Mix	10 µl

4. Start the following PCR reaction program:

- a. Initial Denaturation: 95°C for 3 min.
- b. Denaturation: 95°C for 15 sec
- c. Anneal and Extension: Primer-specific temp. for 60 sec.
- d. Repeat steps b and c for a total of 40 cycles.

5. Analyze quantitative PCR results using software provided with the real-time PCR machine.

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Applications: W—Western IP—Immunoprecipitation IHC—Immunohistochemistry ChIP—Chromatin Immunoprecipitation IF—Immunofluorescence F—Flow cytometry E-P—ELISA-Peptide Species Cross-Reactivity: H—human M—mouse R—rat Hm—hamster Mk—monkey Mi—mink C—chicken Dm—D. melanogaster X—Xenopus Z—zebrafish B—bovine Dg—dog Pg—pig Sc—S. cerevisiae Ce—C. elegans Hr—Horse AII—all species expected Species enclosed in parentheses are predicted to react based on 100% homology